US ERA ARCHIVE DOCUMENT

DP Barcode : D201436 PC Code No : 083301

EEB Out

To: Kathryn Davis

Product Manager 52

DEC 21 1994

Special Review and Reregistration Division (7508W)

From: Anthony F. Maciorowski, Chief

Ecological Effects Branch/EFED (7507C)

Attached, please find the EEB review of...

Reg./File # 083301 Chemical Name

Type Product Product Name

Company Name Purpose

Triazine Joint Venture

Review acute daphnia and oyster shell

studies

Action Code

999

Date Due : 2/15/95

Reviewer Renee Lamb

EEB Guideline/MRID Summary Table:

GDLN NO	MRID NO	CAT	GDLN NO	MRID NO	CAT	GDLN NO	MRID NO	CAT
71-1 (A)			72-2 (A)	431754-01	Y	72-7 (A)		11.
71-1(B)			72-2(B)			72-7 (B)		
71-2 (A)			72-3 (A)			122-1 (A)		
71-2 (B)		Ì.	72-3 (B)	431754-02	Y	122-1 (B)		
71-3			72-3 (C)			122-2		
71-4 (A)			72-3 (D)			123-1 (A)		•
71-4 (B)			72~3 (E)			123-1(B)		
71-5 (A)			72-3 (F)	1		123-2		
71-5(B)			72-4 (A)			123-2		
72-1 (A)		×.	72-4 (B)			123-2		
72-1 (B)			72-5			123-2		1
72-1 (C)		*	72-6			123-2		1
72-1 (C)				-		141-5		

N=Unacceptable (Study was rejected)/Nonconcur

Y=Acceptable (Study satisfied Guideline)/Concur
P=Partial (Study partially fulfilled Guideline but
additional information is needed
S=Supplemental (Study provided useful information but Guideline was

REREG CASE # 3074

DP BARCODE: D201436

SUBMISSION: S462435

CASE: 819287

DATA PACKAGE RECORD

BEAN SHEET

DATE: 04/06/94

Page 1 of 1

\* \* \* CASE/SUBMISSION INFORMATION \* \* \*

ACTION: 627 CORE DATA CASE TYPE: REREGISTRATION

CHEMICALS: 083301 Hexahydro-1,3,5-tris(2-hydroxyethyl)-s-triazine

ID#: 083301

COMPANY:

PRODUCT MANAGER: 52 KATHRYN DAVIS

703-308-8156 ROOM: CS1 3 F 3 4N4

703-308-8523 ROOM: CS1 PM TEAM REVIEWER: BONNIE ADLER DUE OUT DATE: 06/26/94 RECEIVED DATE: 03/28/94

\* \* \* DATA PACKAGE INFORMATION \* \* \*

DP BARCODE: 201436 EXPEDITE: N DATE SENT: 04/06/94 DATE RET.:

CHEMICAL: 083301 Hexahydro-1,3,5-tris(2-hydroxyethyl)-s-triazine

DP TYPE: 999 Miscellaneous Data Package

DATE

04/11/94

LABEL: N CSF: N IN

ASSIGNED TO DIV : EFED

BRAN: EEB

SECT: REVR:

CONTR:

ADMIN DUE DATE: 07/05/94 DATE OUT 11

NEGOT DATE:

PROJ DATE: 2 15/95

\* \* DATA REVIEW INSTRUCTIONS \* \* \*

Please review the following acute tox data for teh chemical grotan;

Acute Toxicity to Daphnia Magna - Flow Through GDLN 72-2a MRID 43175401

GDLN 72-3b Acute Effect on New Shell Growth of Eastern Oyster; MRID 43175402

\* \* \* DATA PACKAGE EVALUATION \* \* \*

No evaluation is written for this data package

\* \* \* ADDITIONAL DATA PACKAGES FOR THIS SUBMISSION \* \* \*

CSF DATE OUT DUE BACK INS BRANCH/SECTION DP BC



# UNITED STATES ENVIRONMENTAL PROTECTION AGENCY WASHINGTON, D.C. 20460

RFG 2 | 1994

OFFICE OF PREVENTION, PESTICIDES AND TOXIC SUBSTANCES

#### **MEMORANDUM**

Subject:

Data review for Triazine (083301)

From:

Anthony F. Maciorowski, Chief

Ecological Effects Branch

Environmental Fate and Effects Division (7507C)

To:

Kathryn Davis, PM 52

Special Review and Reregistration Division (7508W)

EEB has completed the review of the data submitted in support of reregistration of triazine, chemical number 083301. The following is a brief summary of the data reviewed:

1. CITATION: Davis, Jay W. 1994. Triazine: Acute toxicity to the water flea, *Daphnia magna*, under flow-through test conditions. Project No. J9306004d. Performed by Toxikon Environmental Sciences. Submitted by Triazine Joint Venture. EPA MRID No. 431754-01.

**CONCLUSIONS:** This test is scientifically sound and meets the guideline requirements for a flow-through acute toxicity test using *Daphnia magna*. Based on mean measured concentrations, the 48-hour  $EC_{50}$  value is 26.1 mg ai/L (95% C.I. = 19.7 - 34.5 mg ai/L). The NOEC is 7.49 mg ai/L based on mortality at the 13.2 mg ai/L test concentration. Therefore, triazine is classified as slightly toxic to aquatic invertebrates.

**2. CITATION:** Davis, Jay W. 1994. Triazine: Acute effect on new shell growth of the Eastern Oyster, *Crassostrea virginica*. Project No. J9306004h. Prepared by Toxikon Environmental Sciences. Submitted by Triazine Joint Venture. EPA MRID No. 431754-02.

CONCLUSIONS: This study is scientifically sound and meets the guideline requirements for an oyster shell deposition study. The 96-hour EC<sub>50</sub> value is 2.3 mg ai/L (based on mean measured concentrations); the NOEC is 0.87 mg ai/L. Therefore, triazine is classified as moderately toxic to eastern oysters.

If there are any questions contact Renée Costello at 305-5294.

#### DATA EVALUATION RECORD

CHEMICAL: Triazine, Shaughnessey No. 083301

TEST MATERIAL: Technical grade triazine; CAS No. 4719-04-4, Lot No. 0952309, 83.8% active ingredient; a clear yellow viscous liquid.

**STUDY TYPE:** Freshwater Invertebrate Flow-Through Acute Toxicity Test. Species Tested: Daphnia magna.

CITATION: Davis, Jay W. 1994. Triazine: Acute toxicity to the water flea, *Daphnia magna*, under flow-through test conditions. Project No. J9306004d. Performed by Toxikon Environmental Sciences. Submitted by Triazine Joint Venture. EPA MRID No. 431754-01.

#### REVIEWED BY:

Renee Lamb Biologist EFED/EEB Signature, Corol

Date: 6/28/94

#### APPROVED BY:

Ann Stavola
Supervisory biologist, Section 5
EFED/EEB

Signature: allen W. Vaughan

Date: 12.19.94

**CONCLUSIONS:** This test is scientifically sound and meets the guideline requirements for a flow-through acute toxicity test using *Daphnia magna*. Based on mean measured concentrations, the 48-hour EC<sub>50</sub> value is 26.1 mg ai/L (95% C.I. = 19.7 - 34.5 mg ai/L). The NOEC is 7.49 mg ai/L based on mortality at the 13.2 mg ai/L test concentration. Therefore, triazine is classified as slightly toxic to aquatic invertebrates.

#### MATERIALS AND METHODS:

Test Animals: The Daphnia magna (<24 hours old) used in the test were taken from in-house cultures.

Test System: The definitive test was conducted under flow-through conditions in a modified proportional diluter system constructed of glass, silicone adhesive and silicone tubing. A total volume of 53.9  $\mu \rm L$  of test substance was pumped into the chemical mixing chamber each diluter cycle providing a high nominal test concentration of 100 mg ai/L. The test solution was proportionally diluted to provide the 5 lower test concentrations. A dilution water control was also maintained.

The test dilution water was a moderately hard fresh water with a mean hardness of 113 mg/L as CaCO<sub>3</sub>, mean alkalinity os 32 mg/L, and a mean specific conductivity of 658  $\mu$ mhos/cm.

**Dosage:** Six nominal test concentrations - 100, 60.0, 36.0, 21.6, 13.0, and 7.78 mg ai/L and a dilution water control were used.

Design: A test solution of ≈ 220 mL was delivered to each test chamber during every cycle; the total volume was split in halves via a splitter box. Test tanks were 11.3 L glass tanks positioned to provide a maximum depth of 6 cm. Retention chambers were used to prevent neonate floating. Each test container maintained a 450 mL volume of test solution; the diluter cycled at an average rate of 2.9 cycles/hour providing ≈ 17 volume additions every 24 hours.

Ten daphnids were impartially added to each chamber, two chambers per concentration. All test containers were randomly positioned in a single water bath maintained at a temperature of  $20 \pm 1^{\circ}$  C. Fluorescent lighting provided a photoperiod of 16 hours light/8 hours dark, with a 15-minute transition period. The light intensity ranged between 308 and 375 lux.

Survival of daphnids was monitored daily and any dead removed. Abnormalities were also noted. Daphnids were not fed during the test. Test solutions remained clear throughout the study.

Test water quality was monitored daily. Water samples were collected from the controls and all six test concentrations at initiation, day 1, and termination to verify concentrations. Samples were analyzed using HPLC.

**Statistics:** The median effective concentrations (EC $_{50}$ ) and associated 95% confidence intervals (C.I.) were calculated using a computer program (moving average angle, probit, logit, and non-linear interpolation).

REPORTED RESULTS: The diluter functioned properly throughout the test. The mean measured concentrations ranged from 7.49 to 95.2 mg ai.L and from 91 to 102% of nominal. The mean measured concentrations were 7.49, 13.2, 19.7, 34.5, 59.0, and 95.2.

Mortality ranged from 0% at concentrations 7.49 and 19.7 to 100% at concentrations  $\geq$  34.5 mg ai/L. The slope of the concentration response curve could not be determined using the binomial probability method. The 48 hour EC<sub>50</sub> is 26.1 mg ai/L with 95% confidence limits of 19.7 and 34.5 mg ai/L. The NOEC was 19.7 mg ai/L.

Initial alkalinity, hardness and conductivity of the dilution water as measured in the control were 32 mg/L, 130 mg/L, and 828  $\mu$ mhos/cm, respectively. At test termination, they were 32 mg/L,

96 mg/L, and 488  $\mu$ mhos/cm, respectively. During the test, DO remained  $\geq$  7.8 mg/L ( $\geq$  88% saturation). The pH was affected by the presence of triazine with pH increasing with triazine concentration. The pH ranged from 7.7 to 7.8 in the control and from 8.5 to 9.6 in all test concentrations.

STUDY AUTHOR'S CONCLUSIONS/QUALITY ASSURANCE MEASURES: There were no conclusions by the author. A quality assurance statement was included.

#### REVIEWER'S DISCUSSION AND INTERPRETATION OF STUDY RESULTS:

Test Procedure: The study was generally in accordance with SEP and ASTM guidelines.

Statistical Analysis: The reviewer used EPA's Toxanal program to determine the 48-hour  $EC_{50}$  value as 26.1 mg ai/L (95% C.I. = 19.7 - 34.5 mg ai/L). The no-observed-effect concentration (NOEC) can be estimated as 7.49 mg ai/L, based on mortality at the 13.2 mg ai/L concentration The NOEC value is more conservative than the value reported by the study author - the NOEC was reported as 19.7 mg ai/L.

**Discussion/Results:** This test is scientifically sound and meets the guideline requirements for a flow-through acute toxicity test using *Daphnia magna*. Based on mean measured concentrations, the 48-hour EC $_{50}$  value is 26.1 mg ai/L (95% C.I. = 19.7 - 34.5 mg ai/L). The NOEC is 7.49 mg ai/L based on mortality at the 13.2 mg ai/L test concentration.

#### Adequacy of the Study:

- (1) Classification: Core.
- (2) Rationale: N/A.
- (3) Repairability: N/A.

COMPLETION OF ONE-LINER FOR STUDY: June 28, 1994.

NOTE: THERE WAS CONTROL MORTALITY, BUT AT LEAST ONE OF THE LOWER CONCENTRATIONS HAD ZERO MORTALITY. THEREFORE, ABBOTT'S CORRECTION IS NOT APPLICABLE.

lamb	traizine	daphnia
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CONC. NU	MBER	NUMBER	PERCENT	BINOMIAL
EX	POSED	DEAD	DEAD	PROB. (PERCENT)
95.2 2	0	20	100	9.536742E-05
59 2	.0	20	100	9.536742E-05
34.5 2	0	20	100	9.536742E-05
19.7 2	0	0	0	9.536742E-05
13.2 2	0	1	5	2.002716E-03
	0	0	0	9.536742E-05

THE BINOMIAL TEST SHOWS THAT 19.7 AND 34.5 CAN BE USED AS STATISTICALLY SOUND CONSERVATIVE 95 PERCENT CONFIDENCE LIMITS, BECAUSE THE ACTUAL CONFIDENCE LEVEL ASSOCIATED WITH THESE LIMITS IS GREATER THAN 95 PERCENT.

AN APPROXIMATE LC50 FOR THIS SET OF DATA IS 26.0701

WHEN THERE ARE LESS THAN TWO CONCENTRATIONS AT WHICH THE PERCENT DEAD IS BETWEEN 0 AND 100, NEITHER THE MOVING AVERAGE NOR THE PROBIT METHOD CAN GIVE ANY STATISTICALLY SOUND RESULTS.

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#### DATA EVALUATION RECORD

CHEMICAL: Triazine, Shaughnessey No. 083301

TEST MATERIAL: Technical grade triazine, 83.8% active ingredient; a clear yellow viscous liquid.

STUDY TYPE: Mollusc 96-Hour Flow-Through Shell Deposition Study. Species Tested: Eastern oyster (Crassostrea virginica).

CITATION: Davis, Jay W. 1994. Triazine: Acute effect on new shell growth of the Eastern Oyster, Crassostrea virginica. Project No. J9306004h. Prepared by Toxikon Environmental Sciences. Submitted by Triazine Joint Venture. EPA MRID No. 431754-02.

REVIEWED BY:

Renee Lamb Biologist EFED/EEB

APPROVED BY:

Ann Stavola Chief, Section 5 EFED/EEB

Signature:

Date: 6/28/94

Signature: Ollen W. Vaux a

Date: 12.19.94

CONCLUSIONS: This study is scientifically sound and meets the guideline requirements for an oyster shell deposition study. 96-hour  $EC_{50}$  value is 2.3 mg ai/L (based on mean measured concentrations); the NOEC is 0.87 mg ai/L. Therefore, triazine is classified as moderately toxic to eastern oysters.

Test Animals: Eastern oysters (Crassostrea virginica) were obtained from a commercial supplier. During holding, the temperature ranged from 19.0-24.9°C, the salinity ranged from 30-35 ppt. Prior to testing, ≈ 3 - 5 mm of shell growth was removed from the edge of each oyster with a high speed grinder. Actively growing oysters were selected for testing form this group of acclimated oysters and any new shell growth removed immediately before adding to the exposure system. The oysters had an average length of 25-38 mm (mean of 31  $\pm$  3.1 mm) and wet tissue weight of 0.29 to 1.17 g (mean of 0.66  $\pm$  0.29 g).

Oysters were supplementally fed 150 mL of algae per cycle (about 2.5 L per treatment per day).

Test System: The dilution water was natural unfiltered seawater, from the Jupiter River with a salinity of 29 to 33 ppt and a pH of 7.8 to 8.1 during the test.

The definitive test was conducted under flow-through conditions in a modified proportional diluter system constructed of glass, silicone adhesive and silicone tubing. A total volume of 8.41  $\mu L$  of test substance was pumped into the chemical mixing chamber each diluter cycle providing a high nominal test concentration of 15.6 mg ai/L. The test solution was proportionally diluted to provide the 5 lower test concentrations. A dilution water control was also maintained.

**Dosage:** Six nominal concentrations (1.21, 2.02, 3.37, 5.62, 9.36 and 15.6 mg ai/L) and a dilution water control were used in the definitive test.

**Design:** A test solution of  $\approx$  220 mL was delivered to each test chamber during every cycle. Test tanks were 11.3 L glass tanks positioned to provide a maximum depth of 6 cm and a constant water volume of 5.4 L. The diluter cycled at an average rate of 5.5 cycles/hour providing  $\approx$  5.4 volume additions every 24 hours.

Twenty oysters were impartially selected and distributed to each aquarium for a total of 20 oysters per concentration. Oysters were placed equidistantly from one another and facing the incoming flow of water with cupped valves resting on the bottom. Loading was calculated to be  $\approx 0.45$  g of oyster tissue per liter of test solution passing through the test container every 24 hours. Gentle aeration was added to all test containers throughout the test. All tanks were cleaned daily. All test containers were positioned randomly in a single water bath to maintain target temperature of 21  $\pm$  1°. Fluorescent lighting provided a photoperiod of 16 hours light/8 hours dark, with a 15-minute transition period. The light intensity ranged between 317 and 367 lux.

Observations of mortality were made daily. Test water quality were monitored daily during the test. Water samples were collected from the controls and all six test concentrations at initiation, day 1, day 3 and termination to verify concentrations. Samples were analyzed using HPLC.

Statistics: Differences in growth between exposed and control oysters were determined by ANOVA and Dunnett's. The 96-hour EC $_{50}$  value and 95% confidence interval were determined by a computer program (moving average angle, probit, logit, and non-linear interpolation). The method selected for reporting the test results was determined by the characteristics of the data, ie, the presence or absence of 0% and 100% effect and the number of concentrations in which effects between 0 and 100% occurred.

REPORTED RESULTS: The diluter functioned properly throughout the test, except for day 2. The diluter malfunctioned due tot he unfiltered saltwater being diverted to another laboratory source. This diversion caused the diluter to malfunction of the fill

phase of the cycle which also drained the supplementally fed algae source. The problem was corrected and the algae container refilled within 20 minutes of the malfunction.

Chemistry results varied during the test with the best recoveries after test chambers were cleaned; concentrations were lower prior to cleaning. This variability is attributed to indigenous microorganisms from the unfiltered saltwater. (Some biocides or their degradates serve as nutrients stimulating bacterial growth). Following cleaning of the tanks on day 3, chemical recoveries increased over day 1 results supporting this explanation.

DO levels in all treatment levels were higher on day 3 after cleaning on day 2 when the tanks were not cleaned indicating a microbial growth as well. The exception to this pattern was the highest nominal concentration in which DO levels progressively decreased during the test, with only slight differences between day 2 and day 3, indicating that microbial populations at this level may have grown to the extent that simple brushing and rinsing did not sufficiently reduce microbial populations.

Mean measured concentrations were 0.37, 0.87, 1.13, 1.81, 3.16, and 6.52 mg ai/L. These values represent 90 - 112% of nominal concentrations. To demonstrate that test solutions were delivered properly and to confirm that the lower measured concentrations detected in the test chambers during the test resulted from increased bacterial action, the highest test concentration was measured at each sampling time. concentration in the highest test concentration was 10.0, 9.60, 10.1, and 10.7 mg ai/L on days 0, 1, 3, and 4, respectively. mean measured concentration for this level as measured from the delivery tube was 65% of the nominal concentration. At test termination, the measured concentration for 9.36 mg ai/L was less than detection and the delivery tube volume was measured and determined to be 5.08 mg ai/L or 54% of nominal. It is apparent that degradation occurred rapidly in the test system and was responsible for the low measured concentrations.

There were no mortalities during the test. After 96 hours of exposure to triazine, mean new shell growth ranged from 0.27 mm at 6.52 mg ai/L to 2.22 mm at 0.37 mg ai/L. Mean new shell growth in the control was 2.03 mm. The percentage decrease in new shell growth of triazine exposed oysters as compared to the control ranged from 22% at 3.16 mg ai/L to 87% at 6.52 mg ai/L; increases in new shell growth were measured at 0.37 and 0.87 mg ai/L. Growth was statistically reduced from that measured for the control oysters at 1.81 and 6.52 mg ai/L test concentrations.

The 96-hour EC $^{50}$  was 2.31 mg ai/L with 95% confidence limits of 1.13 and 6.52 mg ai/L. The NOEC was 1.13 mg ai/L.

During the test, salinity was 29-33 ppt. The DO ranged from 6.9 to 7.4 mg/L (97 > 100% saturation) at test initiation and remained  $\geq$  5.2 mg/L ( $\geq$  72% saturation) for all control and test solutions. The pH values were 7.8 to 8.1. The test temperature range was 20 to 22.4°C (mean of 21.7  $\pm$  0.6).

STUDY AUTHOR'S CONCLUSIONS/QUALITY ASSURANCE MEASURES:
There were no conclusions by the author. Good Laboratory Practice Compliance and Quality Assurance Statements were included in the report indicating compliance to with EPA Good Laboratory Practice Standards.

#### REVIEWER'S DISCUSSION AND INTERPRETATION OF STUDY RESULTS:

**Test Procedure:** The test procedures were generally in accordance with the SEP. The drop in DO and the low measured concentrations did not affect the results of the study.

Statistical Analysis: The reviewer used EPA's Toxanal program and shell deposition data to determine the 96-hour EC<sub>50</sub> value as 2.3 mg ai/L mean measured concentration using the moving average method. The NOEC was determined to be 0.87 mg ai/L using the raw shell deposition data, and William's test. This NOEC is more conservative than the study author's value of 1.13 mg ai/L.

**Discussion/Results:** This study is scientifically sound and meets the guideline requirements for an oyster shell deposition study. The 96-hour  $EC_{50}$  value is 2.3 mg ai/L (based on mean measured concentrations); the NOEC is 0.87 mg ai/L. Therefore, triazine is classified as moderately toxic to eastern oysters.

### Adequacy of the Study:

- (1) Classification: Core.
- (2) Rationale: N/A.
- (3) Repairability: N/A.

Completion of one liner: June 28, 1994

lamb triazine oyster shell dep

****	********	***	****	****	***	****	* * *
CONC.	NUMBER	NUMBER DEAD		PERCENT DEAD		OMIAL B.(PERCENT	14 (1) 14 (1)
	EXPOSED	DEAD		DEAD	PRO	D. (PEKCENI	1
6.52	100	87		87	0	·	
3.16	100	22	* 1	22	0		
1.81	100	58	•	58	0	$(x_1, \dots, x_n) \in \mathcal{C}_{k}$	
1.13	100	23		23	0		
.87	100	9.		9	0		
.37	100	9		9	0		

BECAUSE THE NUMBER OF ORGANISMS USED WAS SO LARGE, THE 95 PERCENT CONFIDENCE INTERVALS CALCULATED FROM THE BINOMIAL PROBABILITY ARE UNRELIABLE. USE THE INTERVALS CALCULATED BY THE OTHER TESTS.

AN APPROXIMATE LC50 FOR THIS SET OF DATA IS 2.305604

RESULTS CALCULATED USING THE MOVING AVERAGE METHOD SPAN G LC50 95 PERCENT CONFIDENCE LIMITS 3 4.506425E-02 2.779284 2.42744 3.228436

RESULTS CALCULATED USING THE PROBIT METHOD ITERATIONS GOODNESS OF FIT PROBABILITY

0

1.16492 19.01423

A PROBABILITY OF 0 MEANS THAT IT IS LESS THAN 0.001.

SINCE THE PROBABILITY IS LESS THAN 0.05, RESULTS CALCULATED USING THE PROBIT METHOD PROBABLY SHOULD NOT BE USED.

1.800128 SLOPE = 95 PERCENT CONFIDENCE LIMITS =-.1427764 AND 3.743033

LC50 = 2.86802995 PERCENT CONFIDENCE LIMITS = .79747 AND +INFINITY

.5650388 95 PERCENT CONFIDENCE LIMITS = 0 AND 1.439004 \*\*\*\*\*\*\*\*\*\*\*\*\*\* new shell growth for individual oysters

File: b:\grotan\raw.dat Transform: NO TRANSFORM

## SUMMARY STATISTICS ON TRANSFORMED DATA TABLE 1 of 2

GRP	IDENTIFICATION	N	MIN	MAX	MEAN	
1 2 3 4 5 6	control 0.37 0.87 1.13 1.81 3.16 6.52	20 20 20 20 20 20 20 20	1.100 0.000 0.000 0.000 0.000 0.000	3.200 3.800 3.700 3.600 2.500 2.700 1.500	2.025 2.220 2.205 1.560 0.855 1.580 0.265	

## SUMMARY STATISTICS ON TRANSFORMED DATA TABLE 2 of 2

GRP	IDENTIFICATION	VARIANCE	SD	SEM		
1 2 3 4 5 6	control 0.37 0.87 1.13 1.81 3.16 6.52	0.321 0.984 0.917 1.156 0.763 0.540 0.240	0.566 0.992 0.958 1.075 0.873 0.735 0.490	0.127 0.222 0.214 0.240 0.195 0.164 0.110		

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WILLIAMS	TEST	Usoconic	regression	moderi	TABLE' 1	O.F	_

GROUP	IDENTIFICATION	N	ORIGINAL MEAN	TRANSFORMED MEAN	ISOTONIZED MEAN
1 2 3 4 5 6 7	control 0.37 0.87 1.13 1.81 3.16 6.52	20 20 20 20 20 20 20 20	2.025 2.220 2.205 1.560 0.855 1.580 0.265	2.025 2.220 2.205 1.560 0.855 1.580 0.265	2.150 2.150 2.150 1.560 1.218 1.218 0.265

File: b:\grotan\raw.dat Transform: NO TRANSFORM

WILLIAMS TEST	(Isotonic regression model)	TABLE 2 OF 2
IDENTIFICATION	ISOTONIZED CALC. SIG MEAN WILLIAMS P=.05	TABLE DEGREES OF WILLIAMS FREEDOM
control 0.37 0.87 1.13 1.81 3.16 6.52	2.150 2.150 0.471 2.150 0.471 1.560 1.754 * 1.218 3.046 * 1.218 3.046 * 0.265 6.638 *	1.66 k= 1, v=133 1.73 k= 2, v=133 1.75 k= 3, v=133 1.77 k= 4, v=133 1.77 k= 5, v=133 1.78

s = 0.838Note: df used for table values are approximate when v > 20.